Multi-omics Data Integration

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Assistant Professor

June 7th, 2023 | 13:00-15:00 PST

TOG Intermediate Workshop BCCHR Trainee Omics Group (TOG)

Comp Bio lab code



Faculty of Medicine

Centre for

Heart Lung Innovation

UBC and St. Paul's Hospital



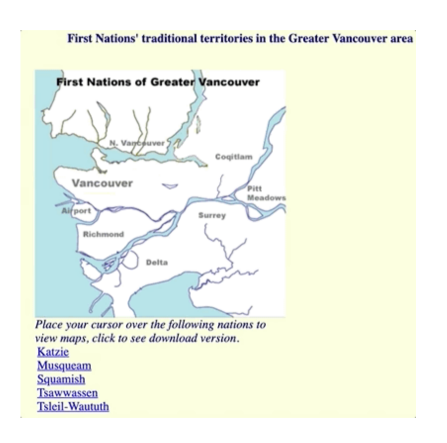
Land acknowledgement

I would like to acknowledge that I work on the traditional, ancestral, and unceded territory of the Coast Salish Peoples, including the territories of the xwməθkwəyəm (Musqueam), Skwxwú7mesh (Squamish), Stó:lō and Səlílwəta?/Selilwitulh (Tsleil-Waututh) Nations.

Traditional: Traditionally used and/or occupied by Musqueam people

Ancestral: Recognizes land that is handed down from generation to generation

Unceded: Refers to land that was not turned over to the Crown (government) by a treaty or other agreement



What are your expectations from today's workshop?





begginer learn new R packages for omics analysis Just exploring

Learning outcomes

By the end of this lecture you will be able to:

- 1. Describe what the *mixOmics* R-library can do.
- 2. Describe when to use which method and for what purpose (exploration, classification, integration).
- 3. Analyze data using mixOmics for various purposes (exploration, classification, integration)

High-dimensional data

- n <<< p (number of observations is much smaller than the number variables)
- data is highly correlated

univariate

p_1 ## 1 -0.646 ## 2 0.544 ## 3 -0.325 ## 4 -0.136 ## 5 -0.630 ## 6 -0.682 ## 7 0.768 ## 8 -0.662 ## 9 0.304 ## 10 0.907

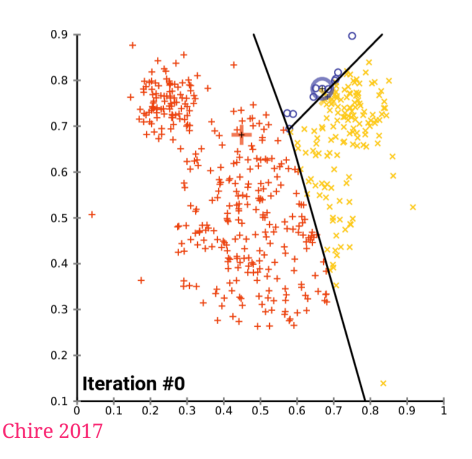
multivariate

```
p_1
                                    p_5
                            p_4
        -1.61000
                         0.5430
                                 0.2980 -0.8360
                                                  0.261
                                                         1.400
-0.7830
         0.06350
                  0.190 -0.2850
                                -0.4670
                                         0.0489 -0.426
                                                        -0.632
-2.6700 -0.77900
                  1.310 -0.2150
                                 0.1790 -1.8700 -0.154
-0.0422 -0.00206 -0.275
                         1.6600
                                 0.9730
                                         0.4910 -0.429
                                                                        0.243
                                                        -0.526
0.0928
         0.36200 -0.539
                         0.0467 -0.2250 -0.2300 -0.924
                                                                        0.891
2.5300
         0.14500 -1.260
                         0.2100
                                 1.0200
                                         1.5700
                                                  0.188
                                                         2.510
-0.0835 -2.05000 -1.050 -0.2660
                                  0.0563
                                         1.1100
                                                  1.200
                                                         0.700
                                                                0.664 -1.460
0.8230
         0.19100 -0.476 1.9300
                                 1.6000 -0.3990 -1.240 -0.189
                                                               -0.496
-1.0000 -0.09970
                  1.630 -0.0202
                                 0.8980
                                         0.4850 -0.549
                                                         0.496 -0.889 -1.440
0.7940 -0.69400
                  0.424 -1.3500 -0.3100 -0.0647 -0.297
                                                         0.684 - 0.908
```

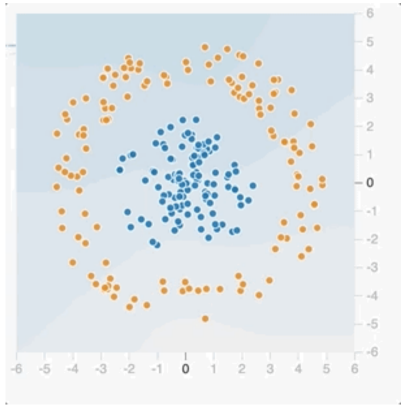


What can you do with high-dimensional data?

Unsupervised (clustering)



Supervised (regression/classification)



Tensorflow playground

mix0mics.org

- initiative started and maintained by Prof Kim-Anh Lê Cao
- R-library with 19 methods for high-dimensional data (exploratory analyses, classification, regression, data integration, meta-analysis)

Lab head: A/Prof Kim-Anh Lê Cao

NHMRC Career Development Fellow

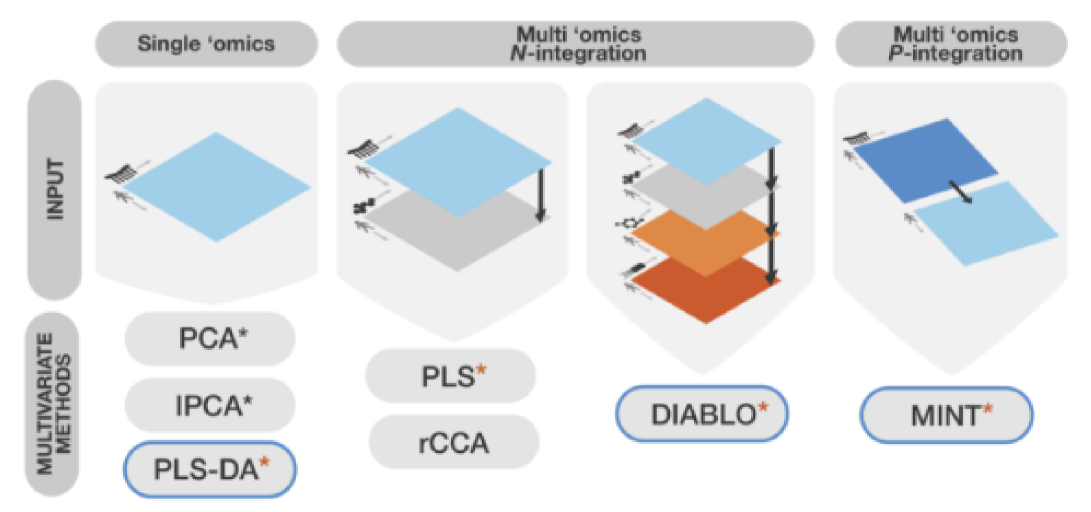
Melbourne Integrative Genomics (MIG) & School
of Mathematics and Statistics

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Melbourne | Parkville VIC 3010

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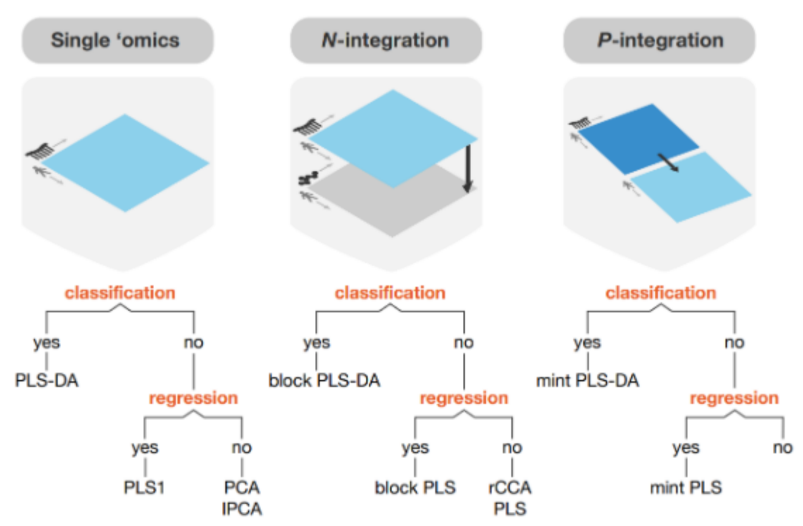


What does mixOmics offer? methods...

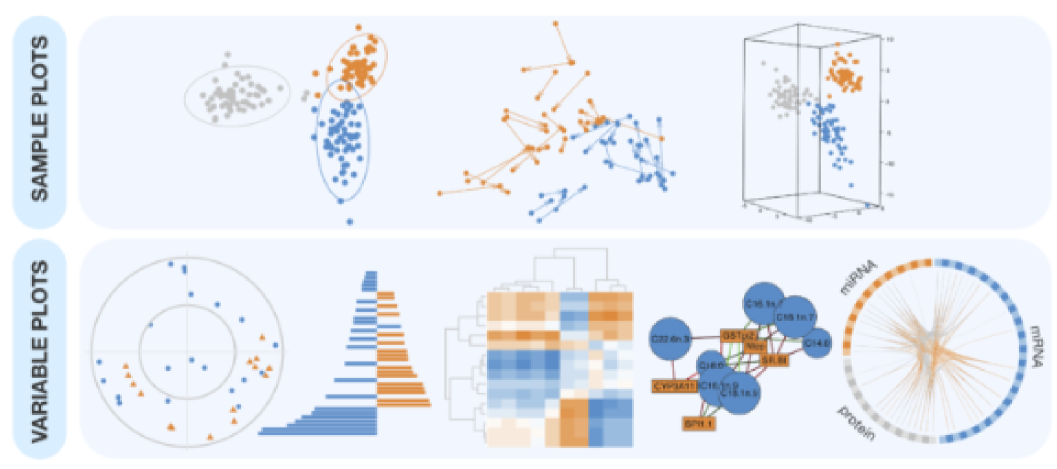


mixOmics.org | *variable selection

What does mixOmics offer? when to use these methods...



What does mix0mics offer? graphics...



mixOmics.org

Getting started with mix0mics

- 1. Download R
- 2. Download RStudio
- 3. install mixOmics

install mixOmics

```
if (!require("BiocManager", quietly = TRUE))
   install.packages("BiocManager")

BiocManager::install("mixOmics")
```

load vignette

```
browseVignettes("mixOmics")
```

Dataset used in this talk

Breast Cancer multi omics data from TCGA

This data set is a small subset of the full data set from The Cancer Genome Atlas that can be analysed with the DIABLO framework. It contains the expression or abundance of three matching omics data sets: mRNA, miRNA and proteomics for 150 breast cancer samples (Basal, Her2, Luminal A) in the training set, and 70 samples in the test set. The test set is missing the proteomics data set.

```
library(mixOmics)
data(breast.TCGA)
lapply(breast.TCGA$data.train, dim)
```

```
## $mirna
## [1] 150 184
##
## $mrna
## [1] 150 200
##
## $protein
## [1] 150 142
##
## $subtype
## NULL
```

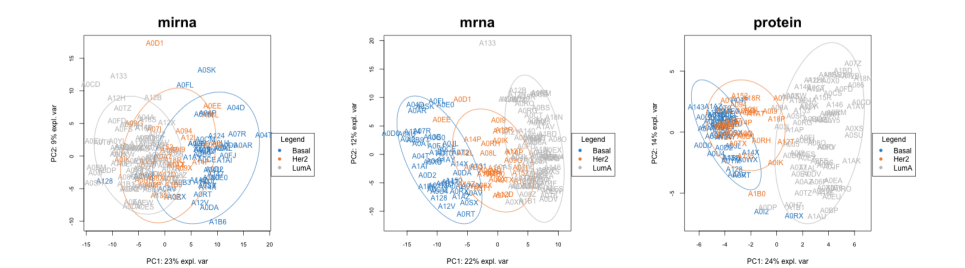
breast cancer subtypes

```
addmargins(table(breast.TCGA$data.train$subtype))
##
## Basal Her2 LumA Sum
## 45 30 75 150
```

Types of analyses covered:

| Analysis | Methods | Functions | Input | Output |
|---------------------------|---------|----------------------------------------------------------|-------------|--------|
| Exploratory data analysis | PCA | pca() plotIndiv() | X | |
| Discriminant analysis | sPLSDA | splsda() tune(), perf() plotIndiv(), plotVar() | X | Y |
| Data integration analysis | DIABLO | block.splsda() tune(), perf() plotDiablo(), circosPlot() | $X_1,, X_J$ | Y |

Exploratory data analysis using PCA



Discriminant analysis using sPLSDA

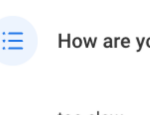
- based on the eda it seems **mrna** is better at separating classes than **mirna**, lets test this.
- this may or may not be true since we peeked at the data (need to test model with another dataset)

• mrna

```
## $overall
          max.dist_centroids.dist_mahalanobis.dist
## comp1 0.22133333
                         0.1760000
                                          0.1760000
## comp2 0.09333333
                         0.1093333
                                          0.1293333
## $BER
          max.dist centroids.dist mahalanobis.dist
## comp1 0.3499259
                        0.1948148
                                         0.1948148
## comp2 0.1191111
                        0.1200000
                                         0.1411852
```

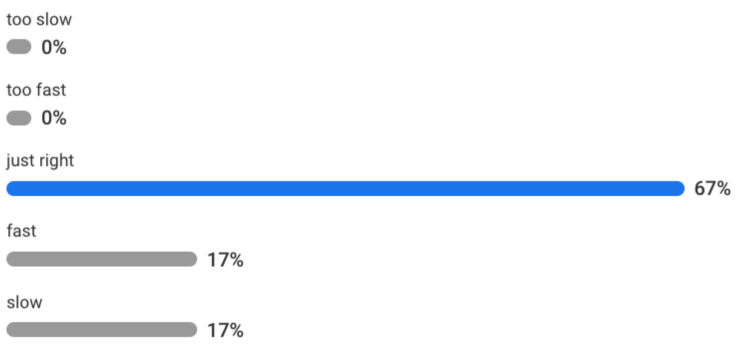
• mirna

```
## $overall
          max.dist_centroids.dist_mahalanobis.dist
                        0.2933333
## comp1 0.2520000
                                         0.2933333
## comp2 0.2133333
                        0.2386667
                                         0.2240000
## $BER
          max.dist centroids.dist mahalanobis.dist
## comp1 0.3774815
                        0.3120000
                                         0.3120000
## comp2 0.2986667
                        0.2708148
                                         0.2682963
```



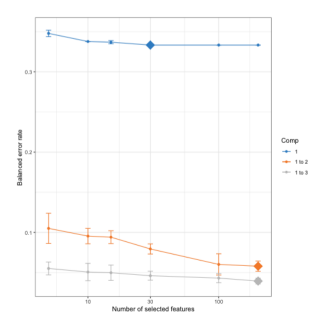
How are you finding the pace of this course?

6 8

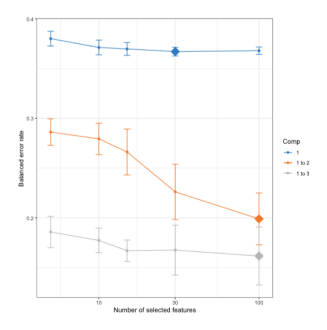


How to select *ncomp* and *keepX*? use a grid of values

• mrna



• mirna

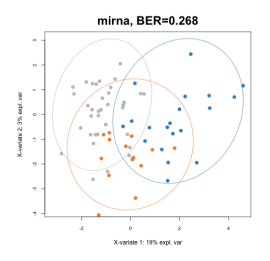


Test sPLSDA models using data from other observations (patients)

• mrna

mrna, BER=0.16

• mirna

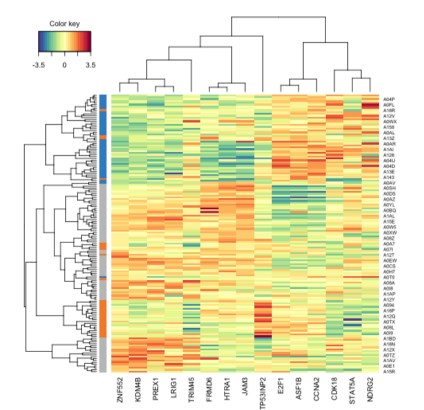


Model interpretation

• determine variables with most importance in mrna model

```
value.var
##
## ZNF552
            -0.75801237
## KDM4B
            -0.58296361
## PREX1
            -0.20979766
## LRIG1
            -0.17185790
## CCNA2
             0.10963799
## CDK18
             0.69045321
## TP53INP2
            -0.68042479
## NDRG2
             0.22678162
## STAT5A
             0.07254351
## TRIM45
             0.06003335
## JAM3
             0.72727214
## E2F1
            -0.53439117
## FRMD6
             0.33920140
## ASF1B
            -0.23142418
## HTRA1
             0.12994836
```

```
cim(mrna_model,
    row.sideColors = mixOmics::color.mixo(as.numeric(b))
```



DIABLO: an integrative classification method for multi-omics data

Design matters!

```
## mrna mirna protein

## mrna 0 1 1

## mirna 1 0 1

## protein 1 1 0
```

```
# set number of component per data set
ncomp = 3
test.keepX = list(mrna = c(10, 30), mirna = c(15, 25),

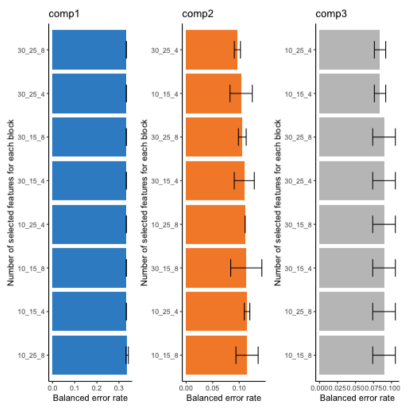
## setup cluster - use SnowParam() on Widnows
BPPARAM <- BiocParallel::MulticoreParam(workers = para
tune <- tune.block.splsda(
    X = data,
    Y = breast.TCGA$data.train$subtype,
    ncomp = ncomp,
    test.keepX = test.keepX,
    design = design,
    nrepeat = 2,
    BPPARAM = BPPARAM
)</pre>
```

```
## Design matrix has changed to include Y; each block will be
## linked to Y.

##
## You have provided a sequence of keepX of length: 2 for block
## This results in 8 models being fitted for each component and
```

Finding the optimal DIABLO model

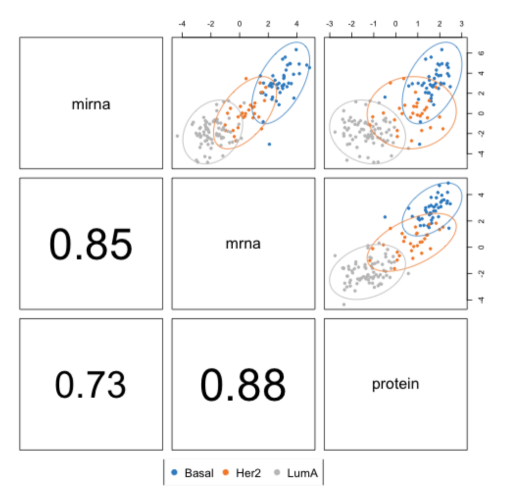




tune\$choice.keepX

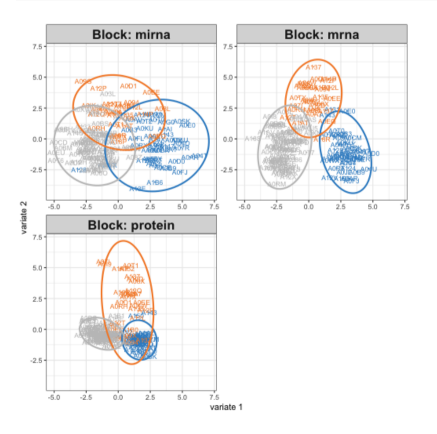
```
## $mrna
## [1] 10 30 10
##
## $mirna
## [1] 15 25 15
##
## $protein
## [1] 4 4 4
```

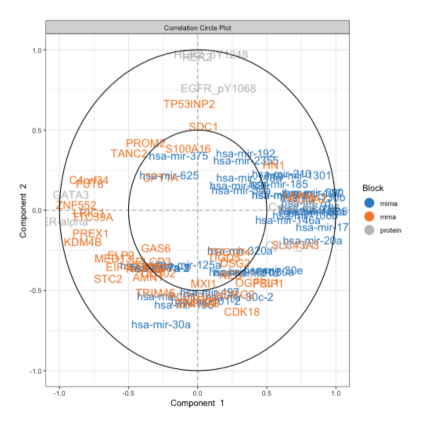
DIABLO model



DIABLO: Sample and variable plots

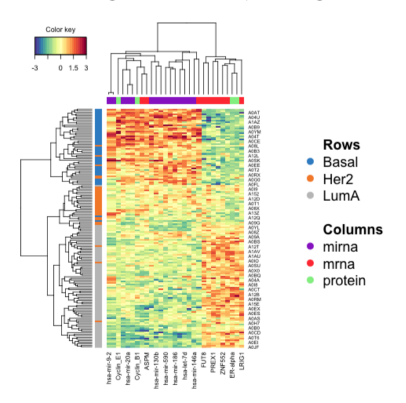
plotIndiv(diablo, ellipse = TRUE)

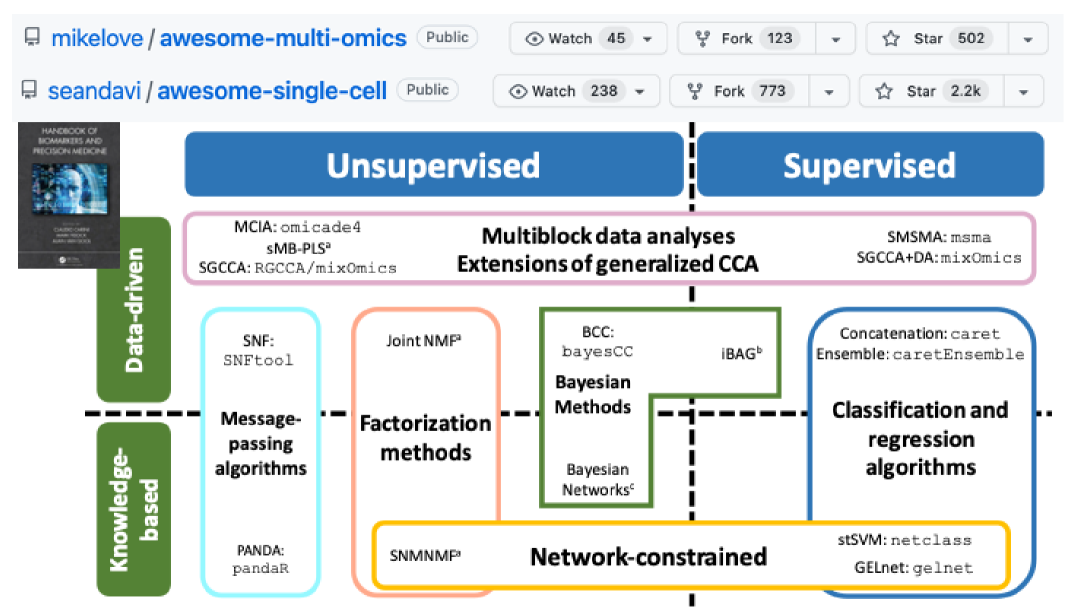




DIABLO

##
trimming values to [-3, 3] range for cim visualisation. See 'trim' arg in ?cimDiablo





Singh A et al., Handbook of Biomarkers and Precision Medicine CRC Press 2019:596



Department of
Anesthesiology, Pharmacology
& Therapeutics
Faculty of Medicine





THANK YOU!

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